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NEWS EXPRESS February 1 CURRENT WINDOWS VERSION IS V6.0d,
CURRENT MACINTOSH VERSION IS V6.0a(ENG) AND V6.0Ja(JP),
AND CURRENT DISCOVER FILE IS DATED 05 FEBRUARY 2002
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NEWS PHONE Direct Dial and Telecommunication Network Access to STN
NEWS WWW CAS World Wide Web Site (general information)

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FILE 'HOME' ENTERED AT 11:07:21 ON 03 OCT 2002

=> file reg
COST IN U.S. DOLLARS

SINCE FILE TOTAL
ENTRY SESSION
0.21 0.21

FULL ESTIMATED COST

FILE 'REGISTRY' ENTERED AT 11:07:32 ON 03 OCT 2002 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2002 American Chemical Society (ACS)

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STRUCTURE FILE UPDATES: 1 OCT 2002 HIGHEST RN 457857-22-6 DICTIONARY FILE UPDATES: 1 OCT 2002 HIGHEST RN 457857-22-6

TSCA INFORMATION NOW CURRENT THROUGH MAY 20, 2002

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Experimental and calculated property data are now available. See HELP PROPERTIES for more information. See STNote 27, Searching Properties in the CAS Registry File, for complete details: http://www.cas.org/ONLINE/STN/STNOTES/stnotes27.pdf

=> ....Testing the current file.... screen

ENTER SCREEN EXPRESSION OR (END):end

=> Uploading C:\STNEXP4\QUERIES\10074014.str

L1 STRUCTURE UPLOADED

=> que L1

L2 QUE L1

=> d 11

L1 HAS NO ANSWERS

L1 STR

G1 Cy,Ak G2 H,M,Cy,Ak

Structure attributes must be viewed using STN Express query preparation.

=> s 11 sss sam

SAMPLE SEARCH INITIATED 11:07:59 FILE 'REGISTRY'

SAMPLE SCREEN SEARCH COMPLETED - 16 TO ITERATE

100.0% PROCESSED 16 ITERATIONS

6 ANSWERS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE \*\*COMPLETE\*\*

BATCH \*\*COMPLETE\*\*

PROJECTED ITERATIONS: 80 TO PROJECTED ANSWERS: 6 TO

L3 6 SEA SSS SAM L1

=> d scan

L3 6 ANSWERS REGISTRY COPYRIGHT 2002 ACS

IN Carbamic acid, [[(2S)-2-amino-3-phenylpropoxy]thioxomethyl]-, ethyl ester, monohydrochloride (9CI)

560

266

MF C13 H18 N2 O3 S . C1 H

Absolute stereochemistry.

HCl

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):5

L3 6 ANSWERS REGISTRY COPYRIGHT 2002 ACS

IN 4,9-Dioxa-2,7-diazaundecanoic acid, 10,10-dimethyl-8-oxo-6-(phenylmethyl)-3-thioxo-, ethyl ester, (6S)- (9CI)

10/074,014

MF C18 H26 N2 O5 S

Absolute stereochemistry.

\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

L3 6 ANSWERS REGISTRY COPYRIGHT 2002 ACS

IN Carbamic acid, (methoxythioxomethyl)-, 2-propenyl ester (9CI)

MF C6 H9 N O3 S

\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

L3 6 ANSWERS REGISTRY COPYRIGHT 2002 ACS

MF C9 H15 N O5 S

\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

L3 6 ANSWERS REGISTRY COPYRIGHT 2002 ACS

IN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 3-butyl 1-ethyl ester,

potassium salt (9CI)

MF C8 H15 N O3 S . K

$$\begin{array}{c|c} & S & O \\ & || & || \\ n\text{-BuO-C-NH-C-OEt} \end{array}$$

• к

L3 6 ANSWERS REGISTRY COPYRIGHT 2002 ACS

IN Uridine, 5'-ester with thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH))
1-ethyl ester (9CI)

MF C13 H17 N3 O8 S

Absolute stereochemistry.

\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

ALL ANSWERS HAVE BEEN SCANNED

=> s ll sss ful

FULL SEARCH INITIATED 11:08:14 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 319 TO ITERATE

100.0% PROCESSED 319 ITERATIONS 104 ANSWERS

SEARCH TIME: 00.00.01

L4 104 SEA SSS FUL L1

=> file caplus

COST IN U.S. DOLLARS
SINCE FILE TOTAL
ENTRY SESSION
140.28 140.49

FILE 'CAPLUS' ENTERED AT 11:08:21 ON 03 OCT 2002

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FILE COVERS 1907 - 3 Oct 2002 VOL 137 ISS 14 FILE LAST UPDATED: 2 Oct 2002 (20021002/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

CAS roles have been modified effective December 16, 2001. Please check your SDI profiles to see if they need to be revised. For information on CAS roles, enter HELP ROLES at an arrow prompt or use the CAS Roles thesaurus (/RL field) in this file.

=> s 14 L5 73 L4

=> s 14/prep

73 L4

2917220 PREP/RL

L6 39 L4/PREP

(L4 (L) PREP/RL)

=> d 16 1-39 bib abs hitstr

- L6 ANSWER 1 OF 39 CAPLUS COPYRIGHT 2002 ACS
- AN 2001:857491 CAPLUS
- DN 135:371451
- TI Improved preparation of high-purity isothiocyanatoformic acid esters as reactants in preparation of pyrrolotriazinones
- IN Matsushita, Akinori
- PA Fuji Photo Film Co., Ltd., Japan
- SO Jpn. Kokai Tokkyo Koho, 15 pp. CODEN: JKXXAF

DT Patent

LA Japanese

FAN.CNT 2

~		_					
	PAT	TENT NO.	KIND	DATE	AP	PLICATION NO.	DATE
ΡI	JΡ	2001328976	A2	20011127	JP	2000-146749	20000518
	US	2002010330	A1	20020124	US	2001-858723	20010517
	US	2002128479	A1	20020912	US	2002-74014	20020214
PRAI	JP	2000-146506	Α	20000518			
	JP	2000-146749	Α	20000518			
	US	2001-858723	A3	20010517			

OS CASREACT 135:371451; MARPAT 135:371451

AB R102CN:C(SR2)OR3 [R1 = (un)substituted alkyl, (un)substituted aryl; R2 = (un)substituted alkyl, (un)substituted aryl, (un)substituted heterocyclyl; R3 = C.gtoreq.3 (un)substituted alkyl, (un)substituted aryl] are prepd. by treatment of ZNCS (Z = Na, K) and R3OH (R3 = same as above) with ClCO2R1 (R1 = same as above) via R102CNHC(:S)OR3 and [R102CNHC(OR3)S]nM (R1, R3 = same as above; M = alkali metal, alk. earth metal, Al, Mg). Thus, ClCO2Et was dropwise added a soln. contg. KNCS and tetrahydrogeraniol at <15.degree. over 1 h, the reaction mixt. stirred at room temp. overnight,

and treated with aq. Ba(OH)2 to give 71% [EtO2CNHC(OR)S]2Ba (R = tetrahydrogeranyl), which was methylated with (MeO)2SO2 in Me2CO to afford the corresponding isothiocyanatoformate with 91% yield and 98% purity.

IT 374540-18-8P 374540-19-9P

RL: IMF (Industrial manufacture); RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (improved prepn. of high-purity isothiocyanatoformic acid esters as reactants in prepn. of pyrrolotriazinones)

RN 374540-18-8 CAPLUS

CN Carbamic acid, [[(3,7-dimethyloctyl)oxy]thioxomethyl]-, ethyl ester, barium salt (9CI) (CA INDEX NAME)

# ●1/2 Ba

RN 374540-19-9 CAPLUS

CN Carbonimidothioic acid, (ethoxycarbonyl)-, O-(3,7-dimethyloctyl) S-methyl ester (9CI) (CA INDEX NAME)

- L6 ANSWER 2 OF 39 CAPLUS COPYRIGHT 2002 ACS
- AN 2001:733456 CAPLUS
- DN 136:53928
- TI Enantioselective total synthesis of batzelladine F: structural revision and stereochemical definition
- AU Cohen, Frederick; Overman, Larry E.
- CS Department of Chemistry, University of California, Irvine, CA, 92697-2025, USA
- SO Journal of the American Chemical Society (2001), 123(43), 10782-10783 CODEN: JACSAT; ISSN: 0002-7863
- PB American Chemical Society
- DT Journal
- LA English

GI

AB The first total synthesis of batzelladine F (I) as the bistrifluoroacetate salt was accomplished in 15 linear steps from two readily available enantiopure .beta.-hydroxy ketones. This enantioselective synthesis revises the structure of batzelladine F and defines its stereochem.

Moreover, the scope of the tethered Biginelli condensation between .beta.-keto ester II as the BF4- salt and guanidine III as the acetate salt has been expanded to include the assembly of complex bisguanidines.

IT 379668-88-9P

RL: RGT (Reagent); SPN (Synthetic preparation); PREP (Preparation)
; RACT (Reactant or reagent)

(asym. total synthesis of batzelladine F via Biginelli condensation, its structure revision and stereochem.)

RN 379668-88-9 CAPLUS

CN Carbamic acid, [methoxy(methylthio)methylene]-, phenylmethyl ester (9CI) (CA INDEX NAME)

RE.CNT 13 THERE ARE 13 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 3 OF 39 CAPLUS COPYRIGHT 2002 ACS

AN 2000:876765 CAPLUS

DN 134:42876

TI Preparation of N-alk(en)oxy(or aryloxy)carbonyl isothiocyanates and their derivatives in the presence of N,N-dialkylarylamine catalysts

IN Kulkarni, Shekhar V.; Desai, Vijay C.

PA Bayer Corporation, USA

SO Eur. Pat. Appl., 12 pp.

CODEN: EPXXDW

DT Patent

LA English

FAN.CNT 2

T.T.	TANGONI Z																	
	PAT	CENT	NO.		KIND DATE				APPLICATION NO.			ο.	DATE					
ΡI	ΕP	1059	289		A.	1	2000	1213		EF	200	00-1	1099	C	20000	0529		
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			IE,	SI,	LT,	LV,	FI,	RO										
	US	6066	754		Α		2000	0523		US	199	99-3	2974	4	19990	0610		
	US	6184	412		В:	1	2001	0206		US	199	99-3	2940	5	19990	0610		
	CA	2310	985		A	Ą	2000	1210		C.P	200	00-2	3109	35	20000	0605		
PRAI	US	1999	-329	405	Α		1999	0610										
	US	1999	-329	744	Α		1999	0610										

OS MARPAT 134:42876

N-alk(en)oxy(or aryloxy)carbonyl isothiocyanates and their derivs. are prepd. by reacting haloformates XCOOR1 (R1 = C1-8 alkyl, C2-4 alkenyl, C6-10 aryl; X = halogen atom; e.g., Me chloroformate) with thiocyanates MSCN (M = alkali metal, alk. earth metal, lead, NH4; e.g., NaSCN) in the presence of catalytic amt. of N,N-dialk(en)ylarylamines (e.g., N,N-dimethylaniline) in aq. solvents or org. solvents to form N,N-dialk(en)ylarylamines (e.g., N-methoxycarbonyl isothiocyanate), and optionally reacting the N,N-dialk(en)ylarylamines with R4YH (R4 = C1-10 alkyl, C6-10 aryl, C1-8 alkoxy; Y = O, S, NR5; R5 = H, R4; e.g., methanol) to form N-alk(en)oxy(or aryloxy)carbonyl isothiocyanate derivs. (e.g., N-methoxycarbonyl-O-methylthionocarbamate) in high yield and purity.

IT 39142-28-4P 39142-31-9P

RL: IMF (Industrial manufacture); **PREP (Preparation)** (prepn. of N-alk(en)oxy(or aryloxy)carbonyl isothiocyanates and their derivs. in the presence of N,N-dialkylarylamine catalysts)

RN 39142-28-4 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), dimethyl ester (9CI) (CA INDEX NAME)

RN 39142-31-9 CAPLUS

RE.CNT 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 4 OF 39 CAPLUS COPYRIGHT 2002 ACS AN 2000:553548 CAPLUS

DN 133:150360

TI Preparation of 2-amino-2-phenylethyl thiocarbamates and analogs as nervous system agents

IN Choi, Yong Moon; Kim, Yong Kil

PA SK Corporation, S. Korea

SO PCT Int. Appl., 75 pp.

CODEN: PIXXD2

DT Patent

LA English

FAN.CNT 1

C. TATA * (	C14 T	Τ.																
	PAI	CENT :	NO.		KI	ND	DATE	DATE AF				CATI	ON NC	Э.	DATE			
PΙ			91	A1 20000810			WC	19	99-KI	R59		19990205						
		W: AU, CA RW: AT, BE		CA,	CN,	JP,	•											
		RW:	AT,	BE,	CH,	CY,	DE,	DK,	ES,	FΙ,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,
			PT,	SE														
	ΑU	9921	890		A	1	2000	0825		JA	J 19	99-2	1890		1999	0205		
	ΕP	1149	076		A	1	2001	1031		EI	2 19	99-90	0198	5	1999	0205		
		R:	BE.	CH.	DE.	ES.	FR.	GB.	IT.	LI.	NL.	SE						

R: BE, CH, DE, ES, FR, GB, PRAI WO 1999-KR59 A 19990205

OS MARPAT 133:150360

AB R(CH2)1CH[(CH2)nNR3R4](CH2)mOCSNR1R2 [I; R = (un)substituted Ph; R1-R4 = H, (cyclo)alkyl, aryl; NR1R2,NR3R4 = heterocyclyl; l,n = 0 or 1; m = 1 or 2] were prepd. Thus, Me3CO2CNHCHPhCH2OH was treated with NaH/CS2 and the product deprotected to give H2NCHPhCH2OCSNH2. Data for biol. activity of I were given.

IT 235439-25-5P 235439-26-6P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of 2-amino-2-phenylethyl thiocarbamates and analogs as nervous system agents)

RN 235439-25-5 CAPLUS

CN Carbamic acid, [[(2R)-2-amino-3-phenylpropoxy]thioxomethyl]-, ethyl ester, monohydrochloride (9CI) (CA INDEX NAME)

Absolute stereochemistry.

# HCl

RN 235439-26-6 CAPLUS

CN Carbamic acid, [[(2S)-2-amino-3-phenylpropoxy]thioxomethyl]-, ethyl ester, monohydrochloride (9CI) (CA INDEX NAME)

Absolute stereochemistry.

# HCl

# IT 235439-48-2P 235439-49-3P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP

(Preparation); RACT (Reactant or reagent)

(prepn. of 2-amino-2-phenylethyl thiocarbamates and analogs as nervous system agents)

RN 235439-48-2 CAPLUS

CN 4,9-Dioxa-2,7-diazaundecanoic acid, 10,10-dimethyl-8-oxo-6-(phenylmethyl)-3-thioxo-, ethyl ester, (6R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 235439-49-3 CAPLUS

CN 4,9-Dioxa-2,7-diazaundecanoic acid, 10,10-dimethyl-8-oxo-6-(phenylmethyl)-3-thioxo-, ethyl ester, (6S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RE.CNT 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 5 OF 39 CAPLUS COPYRIGHT 2002 ACS

AN 2000:344129 CAPLUS

DN 132:321675

TI Process for manufacturing N-alkoxy(or aryloxy)carbonyl isothiocyanate derivatives using N,N-dialkylarylamines as catalysts

IN Kulkarni, Shekhar V.

PA Bayer Corporation, USA

SO U.S., 5 pp. CODEN: USXXAM

DT Patent

LA English

FAN.CNT 2

L'UTA'	OIA I	2																
	PA?	<b>LENT</b>	NO.		KIND		DATE			AP:	PLIC	ATI	ои и	Ю.	DATE			
		<del>-</del>																
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	EΡ	1059	289		A.	1.	2000	1213		EP	200	0-1	1099	0	2000	0529		
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			ΙE,	SI,	LT,	LV,	FI,	RO										
	CA	2310	984		A	Ą	2000	1210		CA	200	0-2	3109	84	2000	0605		
	BR	2000	00259	99	A		2001	0102		BR	200	0-2	599		2000	0608		
	CN	1277	190		Α		2000	1220		CN	200	0-1	1808	5	2000	0609		
	JP	2001	0265	76	$\mathbf{A}_{2}^{\prime}$	2	2001	0130		JP	200	0-1	7366	8	2000	0609		
PRAI	US	1999	-329	405	Α		1999	0610										
	US	1999	-329	744	Α		1999	0610										

OS CASREACT 132:321675; MARPAT 132:321675

AB N-alkoxy(or aryloxy)carbonyl isothiocyanate derivs. R102CNHC(:S)YR4 [R1 = C1-8 alkyl, C2-4 alkenyl, C6-10 aryl; R4 = C1-10 alkyl, C6-10 aryl, C1-8 alkoxy; Y = O, S, NR5; R5 = H, R4] (e.g., N-methoxycarbonyl-O-Me thionocarbamate) are prepd. by reacting a haloformate ester XCO2R1 (X = halogen) (e.g., Me chloroformate) with a thiocyanate MSCN (M = alkali metal, alk. earth metal, NH4) (e.g., sodium thiocyanate) in the presence of an org. solvent (e.g., MIBK) and a catalytic amt. of an 'N,N-dialkylarylamine (e.g., N,N-dimethylaniline) to produce an N-alkoxy(or aryloxy)carbonyl isothiocyanate intermediate S:C:NCO2R1 (e.g., N-methoxycarbonyl isothiocyanate) which then undergoes an addn. reaction with an alc., mercaptan, or amine R4YH (e.g., methanol) to give the N-alkoxy(or aryloxy)carbonyl isothiocyanate deriv. in high yield and purity.

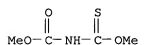
# IT 39142-28-4P

RL: SPN (Synthetic preparation); PREP (Preparation)

(process for manufg. N-alkoxy(or aryloxy)carbonyl isothiocyanate derivs. using N,N-dialkylarylamines as catalysts)

RN 39142-28-4 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), dimethyl ester (9CI) (CA INDEX NAME)



RE.CNT 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 6 OF 39 CAPLUS COPYRIGHT 2002 ACS

AN 1999:505668 CAPLUS

DN 131:144421

TI Preparation of aminoalkyl thiocarbamates as nervous system agents

IN Choi, Yong Moon; Kim, Yong Kil

PA Yukong Limited, S. Korea

SO U.S., 23 pp. CODEN: USXXAM

DT Patent LA English

FAN.CNT 1

PΙ

PATENT NO. KIND DATE APPLICATION NO. DATE
US 5935997 A 19990810 US 1998-6528 19980113

OS MARPAT 131:144421

AB Title compds. (enantiomeric) R(CH2)1CH[(CH2)nNR3R4](CH2)mCH2OCSNR1R2 [R = (un)substituted Ph; R1-R4 = H, (cyclo)alkyl, aryl; NR1R2,NR3R4 = heterocyclyl; l,m,n = 0 or 1] were prepd. as nervous system agents (no data). Thus, PhCH2CH(NHCO2CMe3)CH2OH was treated successively with NaH/CS2, MeI, and aq. NH3 and the product deprotected to give PhCH2CH(NH2)CH2OCSNH2.HCl.

IT 235439-25-5P 235439-26-6P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of aminoalkyl thiocarbamates as nervous system agents)

RN 235439-25-5 CAPLUS

CN Carbamic acid, [[(2R)-2-amino-3-phenylpropoxy]thioxomethyl]-, ethyl ester, monohydrochloride (9CI) (CA INDEX NAME)

Absolute stereochemistry.

# HC1

RN 235439-26-6 CAPLUS

CN Carbamic acid, [[(2S)-2-amino-3-phenylpropoxy]thioxomethyl]-, ethyl ester, monohydrochloride (9CI) (CA INDEX NAME)

Absolute stereochemistry.

## HCl

IT 235439-48-2P 235439-49-3P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP

Absolute stereochemistry.

RN 235439-49-3 CAPLUS

CN 4,9-Dioxa-2,7-diazaundecanoic acid, 10,10-dimethyl-8-oxo-6-(phenylmethyl)-3-thioxo-, ethyl ester, (6S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

# RE.CNT 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 7 OF 39 CAPLUS COPYRIGHT 2002 ACS

AN 1999:417398 CAPLUS

DN 131:58831

TI Process for preparing alkoxytriazolinones

IN Conrad, Michael; Lantzsch, Reinhard; Desai, Vijay C.; Kulkarni, Shekhar V.

PA Bayer Corporation, USA; Bayer Aktiengesellschaft

SO U.S., 6 pp. CODEN: USXXAM

DT Patent

LA English

FAN.CNT 1

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	CA 232	0118		A	A	1999	0819		C	A 19	99-2	3201	18	1999	0130		
	WO 994	1243		Α	1	1999	0819		M	0 19	99-E	P616		1999	0130		
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		ΚE,	KG,	KP,	KR,	KZ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	MD,	MG,	MK,	MN,

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              TR, TT, UA, UG, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
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              CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
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     BR 9907834
                               20001024
                                                BR 1999-7834
                                                                   19990130
                         Α
     EP 1054872
                         Α1
                               20001129
                                                EP 1999-908835
                                                                   19990130
                               20020911
     EP 1054872
                         В1
         R: AT, BE, CH, DE, DK, ES, FR, GB, IT, LI, NL
                         Т2
                               20020205
                                                JP 2000-531438
                                                                   19990130
     JP 2002503654
PRAI US 1998-22262
                               19980211
                         Α
     WO 1999-EP616
                         W
                               19990130
     CASREACT 131:58831; MARPAT 131:58831
OS
GΙ
```

AB Alkoxytriazolinones I (R = alkyl, alkenyl, alkynyl, cycloalkyl, cycloalkylalkyl, aryl, arylalkyl) are prepd. by reacting thioimidodicarboxylic diesters R1O2CNHC(S)OR (R as defined; Rl = alkyl, arylalkyl, aryl) with hydrazine, hydrazine hydrate or an acid adduct of hydrazine. The reaction is conducted in the presence of a diluent and, optionally, in the presence of a basic reaction auxiliary, and at temps. between -10.degree. C. and +100.degree. C.

RN 39142-28-4 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), dimethyl ester (9CI) (CA INDEX NAME)

RN 59701-63-2 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 1-methyl 3-propyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & S & O \\ \parallel & \parallel \\ n\text{-PrO-C-NH-C-OMe} \end{array}$$

# RE.CNT 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L6 ANSWER 8 OF 39 CAPLUS COPYRIGHT 2002 ACS
- AN 1999:120546 CAPLUS
- DN 130:209903
- TI Synthesis and transglycosylase-inhibiting properties of a disaccharide analog of moenomycin A lacking substitution at C-4 of unit F
- AU Riedel, Sylvia; Donnerstag, Astrid; Hennig, Lothar; Welzel, Peter; Richter, Joachim; Hobert, Kurt; Muller, Dietrich; Van Heijenoort, Jean
- CS Institut fur Organische Chemie, Universitat Leipzig, Leipzig, D-04103, Germany
- SO Tetrahedron (1999), 55(7), 1921-1936 CODEN: TETRAB; ISSN: 0040-4020
- PB Elsevier Science Ltd.
- DT Journal
- LA English
- AB A disaccharide analog of moenomycin A lacking the OH group in the 4-position of the uronic acid moiety has been synthesized using the Saito deoxygenation reaction as key step. This analog does not inhibit the transglycosylase (PBP), a key enzyme in the biosynthesis of bacterial peptidoglycan. The result demonstrates the importance of this OH group for the binding of disaccharide moenomycin analogs to the enzyme.
- IT 220974-61-8P

RL: SPN (Synthetic preparation); **PREP (Preparation)** (prepn. and transglycosylase-inhibiting properties of a disaccharide analog of moenomycin A)

RN 220974-61-8 CAPLUS

CN .alpha.-D-Galactopyranosiduronamide, 2-propenyl 2-O-[3,4,6-tri-O-acetyl-2-(acetylamino)-2-deoxy-.beta.-D-glucopyranosyl]-, 3[[(pentafluorophenoxy)thioxomethyl]carbamate] (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RE.CNT 21 THERE ARE 21 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 9 OF 39 CAPLUS COPYRIGHT 2002 ACS

AN 1996:716300 CAPLUS

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DN
    125:328310
    Novel carbamate compounds having N-substituted thiocarbamoyl group, useful
ΤI
    as CNS agents, and process for preparing the same
    Choi, Yong Moon; Han, Dong Il; Kim, Hyung Cheol
TN
    Yukong Limited, S. Korea
PA
SO
     PCT Int. Appl., 22 pp.
     CODEN: PIXXD2
DΤ
    Patent
LA
    English
FAN.CNT 1
                    KIND DATE APPLICATION NO. DATE
    PATENT NO.
    _____
    WO 9632378
                           19961017
                                         WO 1996-KR50
                                                           19960410
                     A1
PΙ
        W: CA, CN, JP
        RW: AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE
                           19981006 US 1996-629619
                                                           19960409
    US 5817858
                Α
                                          CA 1996-2217758
                                                           19960410
    CA 2217758
                      AΑ
                           19961017
                     A1
                                          EP 1996-909387
                           19980128
    EP 820440
                                                           19960410
                      B1
                           20010829
    EP 820440
        R: BE, CH, DE, DK, ES, FR, GB, IT, LI, NL, SE
    CN 1181067 A 19980506
                                     CN 1996-193143
                                                           19960410
                     В
                         20010912
    CN 1070851
    CN 1070851 B 20010912

JP 11503446 T2 19990326

ES 2163010 T3 20020116

CN 1335305 A 20020213
                                          JP 1996-530904
                                                           19960410
                                         ES 1996-909387
                                                           19960410
    CN 1335305
                     A 20020213
                                          CN 2001-101607
                                                           20010117
PRAI KR 1995-8309 A 19950410
WO 1996-KR50 W 19960410
os
    MARPAT 125:328310
    3-O-(N-Substituted-thiocarbamoyl)-2-phenyl-1,3-propanediol carbamates
    H2NCO2CH2CHPhCH2OC(S)NR1R2 (I) are disclosed [wherein R1, R2 = H, C1-8
    alkyl, 5- to 7-membered aliph. cyclic radical optionally contg. .ltoreq. 2
    N or O atoms; both R1 and R2 .noteq. H, and total C in R1 and R2 = 1-16;
    or R1 = H and R2 = C1-8 alkoxycarbonyl or (un)substituted Ph]. I are very
     effective for prophylaxis and treatment of central nervous system
    disorders including nervous muscular pain, epilepsy, and cerebral apoplexy
     (no data). For instance, 2-phenyl-1,3-propanediol monocarbamate in THF
    was treated sequentially with NaH, CS2, and MeI to give 78%
    H2NCO2CH2CHPhCH2OC(S)SMe. This intermediate was treated with aq. MeNH2 in
    THF to give 95% title compd. I [R1 = H, R2 = Me].
IT
    183671-26-3P
    RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use);
     BIOL (Biological study); PREP (Preparation); USES (Uses)
        (prepn. of (thiocarbamoyl)phenylpropanediol carbamates as CNS agents)
RN
     183671-26-3 CAPLUS
CN
     Thioimidodicarbonic acid ((HO)C(O)NHC(S)OH), 3-[3-[(aminocarbonyl)oxy]-2-
```

L6 ANSWER 10 OF 39 CAPLUS COPYRIGHT 2002 ACS

phenylpropyl] 1-ethyl ester (9CI) (CA INDEX NAME)

AN 1995:823077 CAPLUS

DN 123:228001

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Preparation of N-benzoyl-4-acyl- or alkoxypiperidines as substance P
TΙ
     receptor antagonists
IN
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Ofner, Silvio; Roggo, Silvio; Schilling, Walter; Veenstra, Siem J.

PA Ciba-Geigy A.-G., Switz.

SO PCT Int. Appl., 71 pp. CODEN: PIXXD2

DT Patent

LΑ English

FAN.	FAN.CNT 1																	
	PAT	CENT 1	NO.		KII	ND	DATE			A.	PPLI	CATI	ои ис	ο.	DATE			
PI	WO	9511	895		A.	1	1995	0504		W	0 19	94-E	P339	4	1994	1014		
		W:	AM,	AU,	BB,	BG,	BR,	BY,	CA,	CN,	CZ,	EE,	FI,	GE,	HU,	JP,	KG,	ΚP,
			KR,	ΚZ,	LK,	LR,	LT,	LV,	MD,	MG,	MN,	NO,	NZ,	PL,	RO,	RU,	SI,	SK,
			ТJ,	TT,	UA,	US,	UZ,	VN										
		RW:	KE,	MW,	SD,	SZ,	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IE,	IT,	LU,
			MC,	ΝL,	PT,	SE,	BF,	ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	ML,	MR,	NE,	SN,
			TD,	ΤG														
	ΑU	9478	561		A.	1	1995	0522		Αl	U 199	94-7	8561		1994	1014		
PRAI	CH	1993-	-322	3			1993	1026										
	WO	1994-	-EP3	394			1994	1014										
OS	MAF	RPAT :	123:2	2280	01													
GT																		

$$F_3C$$
 $R^{1}N$ 
 $R^{2}X^{1}$ 
 $R^{2}X^{1}$ 
 $R^{2}X^{1}$ 
 $R^{2}X^{1}$ 
 $R^{2}X^{1}$ 
 $R^{2}X^{2}$ 
 $R^{2}X^{2}$ 

AB Title compds. [I; R = X[C(:X4)]nX2X3R3; R1 = aryl(oxy)alkyl, heteroarylalkyl, aroyl, etc.; R2 = cycloalkyl, (un)substituted (hetero)aryl; R3 = (un)substituted (hetero)aryl; R3 = alkyl or (un)esterified or -amidated CO2H when X2 = imino and X3 = alkylene; R4 = H, alkyl, aryl; X,X4 = 0 or S; X1 = bond, CH2, CO, etc.; X2 = bond, (alkyl)imino, alkylene; X3 = bond, alkylene; n = 1; N = 0 when X2 = bondalkylene and X3 = bond] were prepd. Thus, EtOCH2N(CO2CH2Ph)CH(CH2Ph)CH2CH :CH2 (prepn. given) was treated with HCO2H and the deprotected product sequentially N- and O-acylated with 3,5-(F3C)C6H3COCl and 3-bromomethylquinoline, resp., to give title compd. II which had IC50 of 7.6x10-4.mu.M against substance P-induced increase in inositol monophosphate content of human astrocytoma cells (U-373 MG) in vitro.

IT 168271-79-2P

> RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of N-benzoyl-4-acyl- or alkoxypiperidines as substance P receptor antagonists)

RN 168271-79-2 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 3-[1-[3,5-bis(trifluoromethyl)benzoyl]-2-(phenylmethyl)-4-piperidinyl] 1-ethyl ester, trans- (9CI) (CA INDEX NAME)

Relative stereochemistry.

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L6 ANSWER 11 OF 39 CAPLUS COPYRIGHT 2002 ACS
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AN 1995:294083 CAPLUS

DN 123:285785

TI Preparation of aromatic amidine derivatives as inhibitors of human blood coagulation factor for treatment and prevention of influenza

IN Ikeuchi, Kyoshi; Takase, Hiroyuki; Murakami, Yoichi

PA Daiichi Seiyaku Co, Japan

SO Jpn. Kokai Tokkyo Koho, 79 pp. CODEN: JKXXAF

DT Patent

LA Japanese

FAN.CNT 1

PΙ

PATENT NO. KIND DATE APPLICATION NO. DATE

JP 06227971 A2 19940816 JP 1993-17536 19930204

OS MARPAT 123:285785

GI For diagram(s), see printed CA Issue.

The title compds. [I; R1 = H, alkoxy; R2 = H, alkyl, alkoxy, CO2H, alkoxycarbonyl, carboxyalkyl, alkoxycarbonylalkyl; R3 = H, CO2H, alkoxycarbonyl, carboxyalkyl, alkoxycarbonylalkyl, carboxyalkoxy, alkoxycarbonylalkoxy; R4 = H, OH, alkyl, alkoxy; A = C1-4 alkylene which may be substituted by 1-2 of hydroxyalkyl, CO2H, alkoxycarbonyl, carboxyalkyl, and alkoxycarbonylalkyl; X = single bond, O, S, CO; Y = 5or 6-membered (un)satd. carbocyclyl or heterocyclyl, NH2, or aminoalkyl each of which may be substituted; ring Z = pyrrole, 1,2-dihydropyrrole, furan, thiofuran, imidazole, oxazole, thiazole, benzene, tetrahydrobenzene, or cyclopentadiene ring] are prepd. Thus, Et 3-(5-cyano-2-benzofuranyl)-2-(4-hydroxyphenyl)propionate was condensed with (2S)-1-tert-butoxycarbonyl-2-pyrrolidinemethanol in the presence of Ph3P and di-Et azodicarboxylate in THF to give ether (II; R = cyano, R5 =Me3CO2C) which was treated with HCl(g) in ethanol and then with NH3 in EtOH to give amidine II.2HCl (R = amidino, R5 = H). Title compd. (III.2HCl) showed IC50 of 5.04 .mu.g/mL against human blood coagulation.

IT 150613-44-8P, p-Nitrobenzyl N-[methoxy(methylthio)methylene]carbam

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(intermediate for prepn. of arom. amidine derivs. as inhibitors of

human blood coagulation factor)

RN 150613-44-8 CAPLUS

CN Carbonimidothioic acid, [[(4-nitrophenyl)methoxy]carbonyl]-, dimethyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} \text{O} & \text{SMe} \\ \parallel & \parallel \\ \text{CH}_2\text{--O-C-N} & \text{C-OMe} \end{array}$$

L6 ANSWER 12 OF 39 CAPLUS COPYRIGHT 2002 ACS

AN 1995:290316 CAPLUS

DN 122:80880

TI Preparation of 3-methoxy-2-phenylacrylate esters as pesticides.

IN Gayer, Herbert; Gerdes, Peter; Dehne, Heinz-Wilhelm

PA Bayer A.-G., Germany

SO Ger. Offen., 12 pp.

CODEN: GWXXBX

DT Patent

LA German

FAN.CNT 1

GI

1141.	PATENT	NO.		KI	ND	DATE			A	PPLI	CATI	ON NO	0.	DATE			
PI	DE 431																
	WO 942	6705		Α	1	1994	1124		W(	0 19	94-E	P141	7	1994	0504		
	W:	AU,	BB,	BG,	BR,	BY,	CA,	CN,	CZ,	FI,	HU,	JP,	KR,	ΚZ,	LK,	NO,	NZ,
		PL,	RO,	RU,	SK,	UA,	US										
	RW	: AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,
		BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	ML,	MR,	NE,	SN,	TD,	TG		
	AU 946	8427		Α	1	1994	1212		A	J 19	94-6	8427		1994	0504		
	EP 699	183		Α	1	1996	0306		E	P 19	94-9	1693	1	1994	0504		
	R:	BE,	CH,	DE,	ES,	FR,	GB,	IT,	LI,	NL							
	CN 112	3543		Α		1996	0529		CI	N 19	94-1	9213	4	1994	0504		
	JP 085	10217		T	2	1996	1029		J:	P 19	94-5	2490	1	1994	0504		
	US 572	8729		Α		1998	0317		U:	5 19	95-5	5336	0	1995	1113		
PRAI	DE 199	3-431	6431			1993	0517										
	WO 199	4-EP1	417			1994	0504										
os	MARPAT	122:	8088	0													

AB Title compds. [I; R = (substituted) alkyl, cycloalkyl, aryl; n = 0, 1], were prepd. Thus, Me 2-(2-hydroxyphenyl)-3-methoxyacrylate in THF at 0.degree. was treated with 3,5-dichlorobenzoyl isothiocyanate and Et3N; the mixt. was stirred 16 h at room temp. to give title compd. II. Several I gave superior activity against Venturia inaequalis on apples.

IT 160156-92-3P

RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of 3-methoxy-2-phenylacrylate esters as pesticides)

RN 160156-92-3 CAPLUS

CN Benzeneacetic acid, .alpha.-(methoxymethylene)-2[[(phenoxycarbonyl)amino]thioxomethoxy]-, methyl ester (9CI) (CA INDEX NAME)

L6 ANSWER 13 OF 39 CAPLUS COPYRIGHT 2002 ACS

AN 1994:629969 CAPLUS

DN 121:229969

TI Deoxygenation of aliphatic alcohols via reduction of new thioxocarbamate derivatives

AU Oba, Makoto; Nishiyama, Kozaburo

CS Department of Material Science and Technology, Tokai University, Numazu, 410-03, Japan

SO Synthesis (1994), (6), 624-8 CODEN: SYNTBF; ISSN: 0039-7881

DT Journal

LA English

OS CASREACT 121:229969

AB N-Acylthioxocarbamates R1CONHC(:S)OR2 (R1 = e.g., Me), obtained by the reaction of alcs. R2OH (e.g., 1- and 2-dodecanol, cyclododecanol, cholest-5-en-3.beta.-ol) with acyl isothiocyanates, were reduced by tributylstannane or triphenylsilane under radical conditions to give deoxygenated products R2H of the corresponding alcs. in good yields. An application to regioselective deuteration using tributyldeuteriostannane is also examd.

IT 158299-73-1P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP

(Preparation); RACT (Reactant or reagent)

(prepn. and redn. of, by tributylstannane under radical conditions)

RN 158299-73-1 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 1-methyl 3-(1-methylundecyl) ester (9CI) (CA INDEX NAME)

0

S

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O- C- NH- C- OMe
Me-CH-(CH<sub>2</sub>)<sub>9</sub>-Me
    ANSWER 14 OF 39 CAPLUS COPYRIGHT 2002 ACS
1.6
    1994:107001 CAPLUS
ΑN
DN
    120:107001
    Heterocyclic and aromatic amidine derivatives and salts thereof
ΤI
    Nagahara, Takayasu; Kanaya, Naoaki; Inamura, Kazue; Yokoyama, Yukio
IN
    Daiichi Pharmaceutical Co., Ltd., Japan
PA
    Eur. Pat. Appl., 94 pp.
    CODEN: EPXXDW
DT
    Patent
LA
    English
FAN.CNT 1
                                   APPLICATION NO. DATE
                 KIND DATE
    PATENT NO.
                                    _____
    _____
    EP 540051
                 A1 19930505
B1 19960403
                                   EP 1992-118705 19921030
PΤ
    EP 540051
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, NL, PT, SE
    ZA 9208276 A 19930506 ZA 1992-8276 19921026
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US 1992-969369 B1 19921030 US 1992-969396 B1 19921030 US 1994-282571 B3 19940729

US 1995-469593 A1 19950606 US 1997-924504 A3 19970905

OS MARPAT 120:107001

GI For diagram(s), see printed CA Issue.

The title compds. I (where the benzeno-Z ring is indolyl, benzimidazolyl, naphthyl, etc.; R = HN:CNH2; R1 = H, alkoxy; R2 = H, alkyl, alkoxy, etc.; R3 = H, carboxyl, etc.; R4 = H, OH, alkyl, alkoxy; A = C1-4 alkylene; X = single bond, O, S, CO; n = 0-4; Y = heterocyclic or cyclic hydrocarbon moiety) useful as anticoagulant agents were prepd. by treating I (R = CN) with R5OH (R5 = alkyl) to give I (R = R5OC:NH) followed by treatment with NH3. Some of the prepd. compds. showed strong anticoagulant activity through their specific anti-FXa activity in comparison with DABE.

IT 51291-79-3P 150613-44-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP

(Preparation); RACT (Reactant or reagent)

(prepn. and reaction of, in prepn. of amidine anticoagulants)

RN 51291-79-3 CAPLUS

CN Carbonimidothioic acid, (ethoxycarbonyl)-, O-ethyl S-methyl ester (9CI) (CA INDEX NAME)

RN 150613-44-8 CAPLUS

L6 ANSWER 15 OF 39 CAPLUS COPYRIGHT 2002 ACS

AN 1993:653473 CAPLUS

DN 119:253473

TI Lubricating oil containing an antiwear-antioxidant and friction-reducing

IN Beltzer, Morton; Habeeb, Jacob Joseph; Colle, Karla Schall

PA Exxon Research and Engineering Co., USA

SO Eur. Pat. Appl., 13 pp.

CODEN: EPXXDW

DT Patent

LA English

FAN.CNT 1

PRAI US 1991-805757 19911212 US 1992-912539 19920713 OS MARPAT 119:253473

GΙ

$$R^3$$
 $HO \longrightarrow CH_2 \longrightarrow R^5 \longrightarrow NH - CH_2 - CH_2 \longrightarrow II$ 

AB A lubricating oil compn. comprises (a) a base oil and (b) an O-alkyl-N-alkoxycarbonylthionocarbamate having the general formula R1OC(:S)NHC(:O)OR2, where R1 is a hindered phenol having the formula I or an aniline moiety of the formula II, R2 = C1-20 alkyl, aryl, alkaryl, arylalkyl groups, or their substituted derivs., R3, R4 = each a C1-12 alkyl group, and R5 = C2-12 alkyl group. Prefered additives are O-(3,5-di-tert-butyl-4-hydroxybenzyl)-N-ethoxycarbonylthionocarbamate and N,N-((bis-2-hydroxyethyl)-4-hexylanilino)ethoxycarbonylthionocarbamate.

# IT 150929-13-8P

RL: PREP (Preparation)

(prepn. of, antifriction-antioxidants-antiwear additive, for lubricating oils)

RN 150929-13-8 CAPLUS

CN Thioimidodicarbonic acid ((HCO2)NH(HCOS)), O-ethyl O-[2-[(4-hexylphenyl)amino]ethyl] ester (9CI) (CA INDEX NAME)

- L6 ANSWER 16 OF 39 CAPLUS COPYRIGHT 2002 ACS
- AN 1993:427477 CAPLUS
- DN 119:27477
- TI The reactions of some alkoxycarbonyl isothiocyanates with alcohols, phenols and amines
- AU Katritzky, Alan R.; Bernard, Marek K.; Long, Qiu He; Xie, Linghong; Malhotra, Nageshwar; Beltzer, Morton
- CS Cent. Heterocycl. Compd., Univ. Florida, Gainesville, FL, 32611-2046, USA
- SO Organic Preparations and Procedures International (1993), 25(1), 83-90 CODEN: OPPIAK; ISSN: 0030-4948
- DT Journal
- LA English
- AB Reactions of isothiocyanates RO2CNCS (R = Et, dodecyl) with alcs., phenols, and amines were studied. Thus, treatment of dodecyloxycarbonyl isothiocyanate with alcs. gave N-alkoxythiocarbonylcarbamate esters and

with N-heterocycles it gave R1C(S)NHCO2C12H25 (R1 = 1,2,4-triazol-1(or 4)-yl, 1- or 2-benzotriazolyl, or 1-imidazolyl) or R2CO2C12H25 (R2 = 1-benzimidazolyl, 1-pyrazolyl). EtO2CNCS reacted with 4,3,5-HO(Me3C)2C6H2R3 (I; R3 = CH2OH, H) to give I [R3 = OC(S)NHCO2Et or C(S)NHCO2Et, resp.].

IT 148204-38-0P 148204-39-1P 148204-40-4P 148204-41-5P 148204-42-6P 148204-53-9P 148204-56-2P

RN 148204-38-0 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 1-dodecyl ester 3-(2-pyridinylmethyl) ester (9CI) (CA INDEX NAME)

RN 148204-39-1 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 1-dodecyl ester 3-(2-methylpropyl) ester (9CI) (CA INDEX NAME)

RN 148204-40-4 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), didodecyl ester (9CI) (CA INDEX NAME)

RN 148204-41-5 CAPLUS

CN Thioimidodicarbonic acid, 1-dodecyl ester 3-(3-pyridinylmethyl) ester (9CI) (CA INDEX NAME)

RN 148204-42-6 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 1-dodecyl ester 3-1H-pyrrol-2-yl ester (9CI) (CA INDEX NAME)

RN 148204-53-9 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 3-[3,5-bis(1,1-dimethylethyl)-4-hydroxyphenyl] 1-ethyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} O & S \\ \parallel & \parallel \\ EtO-C-NH-C-O-CH_2 & Bu-t \\ \hline \\ OH & \\ \hline \\ t-Bu & \end{array}$$

RN 148204-56-2 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 3-dodecyl 1-ethyl ester (9CI) (CA INDEX NAME)

- L6 ANSWER 17 OF 39 CAPLUS COPYRIGHT 2002 ACS
- AN 1991:449530 CAPLUS
- DN 115:49530
- TI A convenient synthesis of 1,2,4-oxadiazolidine-3,5-dione
- AU Renaut, P.; Thomas, D.; Bellamy, F. D.
- CS Lab. Fournier, Cent. Rech., Fontaine les Dijon, F-21121, Fr.
- SO Synthesis (1991), (4), 265-6 CODEN: SYNTBF; ISSN: 0039-7881
- DT Journal
- LA English
- OS CASREACT 115:49530

GΙ

AB Title compd. I was prepd. by condensation of PhCH2OH with EtO2CNCS to give EtO2CNHC(S)OCH2Ph which cyclized with NH2OH to give 3-benzyloxy-1,2,4-

oxadiazol-5(4H)-one which was debenzylated using BBr3.

IT 59965-72-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP.

(Preparation)

(prepn. and cyclocondensation of, with hydroxylamine)

RN 59965-72-9 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 1-ethyl 3-(phenylmethyl)
 ester (9CI) (CA INDEX NAME)

L6 ANSWER 18 OF 39 CAPLUS COPYRIGHT 2002 ACS

AN 1991:122217 CAPLUS

DN 114:122217

TI The Friedel-Crafts reaction of phenols with carbethoxy isothiocyanate

AU Jagodzinski, Tadeusz

CS Dep. Org. Chem., Tech. Univ. Szczecin, Szczecin, 71-065, Pol.

SO Org. Prep. Proced. Int. (1990), 22(6), 755-60 CODEN: OPPIAK; ISSN: 0030-4948

DT Journal

LA English

OS CASREACT 114:122217

The Freidel-Crafts reaction of phenol derivs. with EtO2CNCS (I) was dependent on the homogeneity of the reaction mixt. Thus, the reaction of C6H5OH with I in the presence of AlCl3 in CH2Cl2 or THF/Et3N gave 98% O-alkylated product, i.e., PhOC6H4OC(S)NHCO2Et. The reaction of C6H5OH with I in the presence of AlCl3 in MeNO3 gave 89% C-alkylated product, i.e., 4-HOC6H4C(S)NHCO2Et. The reaction of 2-naphthalenol with I gave 3,4-dihydro-2-oxo-2H-naphth[1,2-e]-1,3-oxazin-4-thione. A reaction mechanism was discussed.

IT 132554-63-3P

RN 132554-63-3 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 1-ethyl 3-phenyl ester (9CI) (CA INDEX NAME)

L6 ANSWER 19 OF 39 CAPLUS COPYRIGHT 2002 ACS

AN 1990:514622 CAPLUS

DN 113:114622

TI Preparation of ethoxycarbonyl isothiocyanate using a pyridine or quinoline catalyst

AU Lewellyn, Morris E.; Wang, Samuel S.; Strydom, Peter J.

CS Chem. Res. Div., American Cyanamid Co., Stamford, CT, 06904, USA

SO J. Org. Chem. (1990), 55(18), 5230-1 CODEN: JOCEAH; ISSN: 0022-3263

DT Journal

LA English

OS CASREACT 113:114622

AB A process for the prepn. of EtO2CNCS (I) from ClCO2Et and NaSCN using pyridine or quinoline as a catalyst in an aq. medium is presented. This process leads to high yields of the desired product with only trace amts. of the thiocyanate being formed. The reactions of nucleophiles with I, prepd. in situ, can be carried out in high yields and purity.

IT 103122-66-3P

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of)

RN 103122-66-3 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 1-ethyl 3-(2-methylpropyl) ester (9CI) (CA INDEX NAME)

L6 ANSWER 20 OF 39 CAPLUS COPYRIGHT 2002 ACS

AN 1988:37225 CAPLUS

DN 108:37225

TI Preparation of iodopropargylurethanes as pesticides

IN Brandes, Wilhelm; Bunnenberg, Rolf; Reinecke, Paul; Paulus, Wilfried; Schmitt, Hans Georg

PA Bayer A.-G., Fed. Rep. Ger.

SO Ger. Offen., 11 pp.

CODEN: GWXXBX

DT Patent

LA German

FAN.CNT 1

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 3607624	A1	19870910	DE 1986-3607624	19860307

OS CASREACT 108:37225

GΙ

PΙ

AB IC.tplbond.CCH2OCONHCSR [I; R = (substituted) arylthio, alkylthio, alkoxy, alkylamino, etc.] are prepd. as pesticides. A mixt. of 50 mmol each of IC.tplbond.CCH2OH and thiadiazinone II THF was stirred at 0-25.degree. for several hours in the presence of Et3N to give 26% I (R = NMePh). I (R = 4-ClC6H4S) proved effective in tests as an algicide, fungicide, and pesticide.

IT 112111-84-9P

RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(prepn. of, as pesticide, fungicide and algicide)

112111-84-9 CAPLUS RN

Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 3-ethyl CN 1-(3-iodo-2-propynyl) ester (9CI) (CA INDEX NAME)

ANSWER 21 OF 39 CAPLUS COPYRIGHT 2002 ACS L6

1987:439240 CAPLUS ΑN

107:39240

Process for the production of isothiocyanate derivatives TI

Fu, Yun-lung; Strydom, Peter J. IN

American Cyanamid Co., USA PA

SO U.S., 6 pp. CODEN: USXXAM

DT Patent

English LΑ

FAN.CNT 1

PATENT NO. KIND DATE APPLICATION NO. DATE

US 4659853 A 19870421 US 1986-821297 19860122 PΤ

Carbonyl isothiocyanate derivs. (i.e. thionocarbamates, thioureas, and dithiocarbamates) were prepd. in high yields in a one-pot process wherein RO2CX (R = C1-8 alkyl, C3-4 alkenyl, C6-10 aryl; X = halo) were treated with MSCN (M = alkali or alk. earth metal, Pb, NH4) and subsequently with R1YH (R1 = C1-10 alkyl, C6-10 aryl, C1-8 alkoxy; Y = O, S, NR2; R2 = H, R1). NaSCN reacted with ClCO2Et in the presence of pyridine to give EtO2CNCS, which was esterified with iso-BuOH to give EtO2CNHC(S)OCH2CHMe2 (85%).

103122-66-3P 103122-67-4P 109202-54-2P IT

109202-55-3P 109202-58-6P

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of)

103122-66-3 CAPLUS RN

Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 1-ethyl 3-(2-methylpropyl) CN ester (9CI) (CA INDEX NAME)

RN 103122-67-4 CAPLUS

Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 1-ethyl 3-pentyl ester CN (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & S & O \\ \parallel & \parallel & \parallel \\ \text{Me- (CH2)}_4 - O - C - NH - C - OEt \end{array}$$

RN 109202-54-2 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 3-(2-methylpropyl) 1-phenyl ester (9CI) (CA INDEX NAME)

RN 109202-55-3 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 3-(2-methylpropyl) 1-(2-propenyl) ester (9CI) (CA INDEX NAME)

RN 109202-58-6 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 1-ethyl 3-hexyl ester (9CI) (CA INDEX NAME)

L6 ANSWER 22 OF 39 CAPLUS COPYRIGHT 2002 ACS

AN 1986:427656 CAPLUS

DN 105:27656

TI Collectors and froth flotation processes for metal sulfide ores

IN Fu, Yun Lung; Wang, Samuel Shan Ning; Nagaraj, Devarayasamudram Ramachandran

PA American Cyanamid Co., USA

SO Brit. UK Pat. Appl., 36 pp. CODEN: BAXXDU

DT Patent

LA English

FAN.CNT 3

ATENT NO.	KIND	DATE	APPLICATION NO.	DATE
В 2163068	A1	19860219	GB 1985-19737	19850806
В 2163068	B2	19880928		
s 4556482	Α	19851203	US 1984-641659	19840817
S 4556483	Α	19851203	US 1984-641660	19840817
s 4584097	Α	19860422	US 1984-641657	19840817
s 4595493	Α	19860617	US 1984-641658	19840817
A 1278111	A1	19901218	CA 1985-488780	19850815
	B 2163068 B 2163068 S 4556482 S 4556483 S 4584097 S 4595493	B 2163068 A1 B 2163068 B2 S 4556482 A S 4556483 A S 4584097 A S 4595493 A	B 2163068 A1 19860219 B 2163068 B2 19880928 S 4556482 A 19851203 S 4556483 A 19851203 S 4584097 A 19860422 S 4595493 A 19860617	B 2163068 A1 19860219 GB 1985-19737 B 2163068 B2 19880928 S 4556482 A 19851203 US 1984-641659 S 4556483 A 19851203 US 1984-641660 S 4584097 A 19860422 US 1984-641657 S 4595493 A 19860617 US 1984-641658

		8506249	Α	19860326		1985-6249	19850816
	US	4657688	Α	19870414	US	1985-806585	19851209
	US	32827	E	19890110	US	1987-79629	19870730
	GB	2193660	A1	19880217	GB	1987-18337	19870803
	GB	2193660	B2	19880928			
PRAI	US	1984-641657		19840817			
	US	1984-641658		19840817			•
	US	1984-641659		19840817			
	US	1984-641660		19840817			
	GB	1985-19737		19850806			
	US	1985-806585		19851209			

Collectors for sulfide minerals suitable for a broad pH range comprise AB hydrocarbyloxycarbonyl thionocarbamate(I) or similar thiourea(II) compds. added at 0.005-0.5 lb/ton ore. Froth flotation at pH <10 (preferably 4-10) decreases lime consumption and permits a selective rejection of pyrite and pyrrhotite. The I compds. are R1OC(:0)N(H)C(S)OR2 having R1 and R2 as hydrocarbyl, alkyl polyether, and/or arom. radicals. optionally substituted with polar halogen, nitrile, or nitro groups, preferably with R1 as C1-6 alkyl or aryl and R2 as C1-8 alkyl. The II compds. are R3OC(:O)N(H)C(S)NR1R2 having R1 as H or R2, esp. H or C1-6 alkyl; R2 as a hydroxycarbyl, hydrocarboxy, or arom. radical, preferably C1-8 alkyl, allyl, alkaryl, or aryl; and R3 as hydrocarbyl, alkyl polyether, or arom. radical, preferably C1-6 alkyl or aryl. Thus, powd. sulfide ore contg. 0.3 Cu and 1.7% pyrite was slurried at natural pH 5.5 for 30% solids, and conditioned for flotation with collector and frother. In tests with o-iso-Pr N-(ethoxycarbonyl) thiocarbamate at 0.054 lb/ton ore the Cu recovery was 90.8% at conc. grade 9.6% and pyrite recovery 67.3%, compared with 73.2, 2.7, and 57.1 resp. for o-iso-Pr N-ethylthiocarbamate.

IT 58902-91-3P 103122-66-3P 103122-67-4P

RL: IMF (Industrial manufacture); PREP (Preparation) (prepn. of, for collectors in froth flotation)

RN 58902-91-3 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 3-ethyl 1-phenyl ester (9CI) (CA INDEX NAME)

RN 103122-66-3 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 1-ethyl 3-(2-methylpropyl)
 ester (9CI) (CA INDEX NAME)

RN 103122-67-4 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 1-ethyl 3-pentyl ester (9CI) (CA INDEX NAME)

L6 ANSWER 23 OF 39 CAPLUS COPYRIGHT 2002 ACS

AN 1985:149700 CAPLUS

DN 102:149700

TI Synthesis and cytostatic activity of 5'-O-substituted nucleosides

AU Garcia-Lopez, M. T.; Fernandez-Resa, P.; De las Heras, F. G.; Mendez-Castrillon, P. P.

CS Inst. Quim. Med., CSIC, Madrid, 28006, Spain

SO An. Quim., Ser. C (1984), 80(2), 168-71 CODEN: AQSBD6; ISSN: 0211-1357

DT Journal

LA Spanish

GΙ

AB Uridine derivs. I and inosine derivs. II (R = H, Ac or R2 = Me2C; R1 = C1CH2CO, ICH2CO, EtO2CNHCS, H2NCO) were prepd. by acylation of I and II (R2 = Me2C, R1 = H) and optional iodine-chlorine exchange and deprotection reactions. The iodoacetylated nucleosides, esp. I (R2 = Me2C, R1 = ICH2CO), showed significant cytostatic activity against HeLa cell cultures.

IT 95578-11-3P 95578-12-4P

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. and cytostatic activity of)

RN 95578-11-3 CAPLUS

CN Inosine, 2',3'-O-(1-methylethylidene)-, 5'-ester with thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)) 1-ethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 95578-12-4 CAPLUS

CN Uridine, 5'-ester with thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)) 1-ethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

IT 95578-10-2P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP

(Preparation)

(prepn., reactions, and cytostatic activity of)

RN 95578-10-2 CAPLUS

CN Uridine, 2',3'-O-(1-methylethylidene)-, 5'-ester with thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)) 1-ethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L6 ANSWER 24 OF 39 CAPLUS COPYRIGHT 2002 ACS

1984:570775 CAPLUS AN

DN 101:170775

2,2-Dichloro-3,3-dimethylcyclopropylmethyl carbamate derivatives as TΙ fungicides

Nihon Tokushu Noyaku Seizo K. K., Japan PA

Jpn. Kokai Tokkyo Koho, 13 pp. SO

CODEN: JKXXAF

DΤ Patent

Japanese LA

FAN.CNT 1

APPLICATION NO. DATE PATENT NO. KIND DATE -----\_\_\_\_ \_\_\_\_\_ 19840612 JP 1982-211239 19821203 JP 59101453 A2 ΡI GΙ

AΒ Twenty title carbamates (I; R, R1 = H, Me; R2 = H, alkyl, aryl; R1R2N = heterocycle; Z = O, S), effective fungicides at 200 mg/m2, were prepd. Thus, a mixt. of 1.7 g II and 1.0 g MeNCO in CH2Cl2 contg. NaOMe was refluxed 12 h to give 1.90 g I (R = R1 = H, R2 = Me, Z = O).

ΙT 92533-76-1P

> RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of)

RN 92533-76-1 CAPLUS

Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 3-[(2,2-dichloro-3,3-CN dimethylcyclopropyl)methyl] 1-ethyl ester (9CI) (CA INDEX NAME)

L6 ANSWER 25 OF 39 CAPLUS COPYRIGHT 2002 ACS

AN 1982:162095 CAPLUS

DN 96:162095

Thioacyl isocyanates. XVI. Ethoxy(thiocarbonyl) isocyanate Goerdeler, Joachim; Schulze, Andreas TI

AU

Inst. Org. Chem. Biochem., Univ. Bonn, Bonn, D-5300/1, Fed. Rep. Ger. Chem. Ber. (1982), 115(3), 1252-5CS

SO CODEN: CHBEAM; ISSN: 0009-2940

DTJournal

German LА

GΙ

AB EtOC(S)NCO (I) was prepd. by treating EtOC(S)NH2 with (COCl)2 in HCCl3. I dimerizes readily to give II, which is a good starting material for the monomer. The I/II ratio was detd. in PhNO2 at 93.degree. and the reactions of I with some nucleophiles are reported.

IT 59386-42-4P

RN 59386-42-4 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 3-ethyl 1-methyl ester (9CI) (CA INDEX NAME)

L6 ANSWER 26 OF 39 CAPLUS COPYRIGHT 2002 ACS

AN 1982:104273 CAPLUS

DN 96:104273

TI Carbonylthiocarbonylamine compounds

IN Jochims, Johannes Christian; Bunnenberg, Rolf

PA Bayer A.-G., Fed. Rep. Ger.

SO Ger. Offen., 26 pp.

CODEN: GWXXBX

DT Patent

LA German

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI GI	DE 3010204	A1	19810924	DE 1980-3010204	19800317

AB Title compds. RCONHCSR1 [R = R2X, R3XCSNH [R2 = alkyl, aryl; R3 = aryl, X

10/074,014

= O, S, NR4 (R4 = H, alkyl, aralkyl, aryl)]; R1 = ZR5 [R5 = alkyl, aralkyl, aryl, Z = O, S, NR6 (R6 = H, alkyl, aralkyl, aryl)]; R, R1 = morpholino, piperidino, piperazino, thiomorpholino]; the triazines I (R7 = aralkyl, aryl) and thiadiazines II (R8 = R4, morpholino, piperidino, piperazino; thiomorpholino) were prepd. Thus, CO(SCN)2, prepd. from ammonium thiocyanate and Cl2CO, was treated with PhCH2NH2 to give 6-(benzylamino)-2,3-dihydro-2-thioxo-4H-1,3,5-thiadiazin-4-one, which was rearranged and the resulting 1-benzyl-1,2,3,4,5,6-hexahydro-2,6-dithioxo-1,3,5-triazin-4-one treated with PhNH2 to give 1-benzyl-7-phenyl-2,6-dithiotriuret. CO(SCN)2 was treated with (Me2CH)2NH to give 1,5-diisopropyl-2-thiobiuret.

IT 39142-28-4P

RN 39142-28-4 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), dimethyl ester (9CI) (CA INDEX NAME)

L6 ANSWER 27 OF 39 CAPLUS COPYRIGHT 2002 ACS

AN 1981:532748 CAPLUS

DN 95:132748

TI Carbonyl diisothiocyanate

AU Bunnenberg, Rolf; Jochims, Johannes C.

CS Fachber. Chem., Univ. Konstanz, Konstanz, D-7750, Fed. Rep. Ger.

SO Chem. Ber. (1981), 114(6), 2075-86 CODEN: CHBEAM; ISSN: 0009-2940

DT Journal

LA German

GI

AB CO(NCS)2 (I) was prepd. in 50-70% yield by treating NH4SCN with COC12 in THF at 3.degree.. I, a very strong electrophile, is sol. in all org. solvents, but reacts explosively with Me2SO. Treating I with H2O or H2S gave II (Z = O, S) or with alcs., mercaptans, or amines gave III (R = OMe, OPh, SEt, Et2N, PhNH, etc.); excess nucleophile cleaved the heterocyclic ring with addn. or substitution of the resulting NCS group. Thus, I reacted with excess NH3 to give H2NCONHCSNH2; treating III (R = NHPh) with PhNH2 or PhCH2NH2 gave PhNHCSNHCONHCSNHPh and PhCH2NHCONHCSNHPh, resp. Thermal rearrangement of III (R = NHPh, NHCH2Ph) gave IV (R2 = Ph, CH2Ph).

IT 39142-28-4P

RN 39142-28-4 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), dimethyl ester (9CI) (CA INDEX NAME)

L6 ANSWER 28 OF 39 CAPLUS COPYRIGHT 2002 ACS

AN 1980:6471 CAPLUS

DN 92:6471

TI Effect of electrophilic reagents on the 3-hydroxy-1,2,4-thiadiazoles

AU Taliani, Laurent; Perronnet, Jacques

CS Cent. Rech., Roussel-Uclaf, Romainville, 93230, Fr.

SO J. Heterocycl. Chem. (1979), 16(5), 961-71

CODEN: JHTCAD; ISSN: 0022-152X

DT Journal

LA French

GΙ

AB Electrophilic reagents may react either with the hydroxyl group in position 3, or with the 2-nitrogen atom of 3-hydroxy-1,2,4-thiadiazoles (I; R = alkoxy, alkylthio, NMe2). Hard electrophiles, such as acid chlorides, substitute on OH, whereas soft electrophiles (isocyanates, acid anhydrides) substitute on N.

IT 59965-65-0P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP

(Preparation)

(prepn. and cyclization of)

RN 59965-65-0 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 3-[(4-chlorophenyl)methyl] 1-ethyl ester (9CI) (CA INDEX NAME)

L6 ANSWER 29 OF 39 CAPLUS COPYRIGHT 2002 ACS

AN 1976:478112 CAPLUS

85:78112 DN

Pesticidal organophosphorus thiazole derivatives TI

Perronnet, Jacques; Taliani, Laurent IN

PΑ Roussel-UCLAF, Fr.

SO Ger. Offen., 35 pp.

CODEN: GWXXBX

DT Patent

German LΑ

FAN.CNT 1						
PA'	TENT NO.	KIND	DATE	APPLICATION NO.	DATE	
PI DE	2541720	A1	19760408	DE 1975-2541720	19750918	
FR	2285397	<b>A</b> 1	19760416	FR 1974-31841	19740920	
US	4020076	A	19770426	US 1975-611710	19750909	
JP	51059859	A2	19760525	JP 1975-112201	19750918	
ES	441054	A1	19770301	ES 1975-441054	19750918	
BE	833618	<b>A</b> 1	19760319	BE 1975-160203	19750919	
DK	7504207	A	19760321	DK 1975-4207	19750919	
DK	138747	С	19790402			
DK	138747	В	19781023			
NL	7511066	Α	19760323	NL 1975-11066	19750919	
BR	7506044	A	19760803	BR 1975-6044	19750919	
CH	602776	A	19780731	CH 1975-12172	19750919	
CA	1056384	<b>A</b> 1	19790612	CA 1975-236072	19750919	
GB	1502890	Α	19780308	GB 1975-38757	19750922	
PRAI FR	1974-31841		19740920			
GI						

- Thiazolyl phosphates I (R = OEt, OBu, cyclohexyloxy, OCH2C6H4Cl-4, SEt, AB OCH2Ph, SCH2C6H4Cl-4, OCH2C6H4Me-4; R1 = Me, Et; X = O, S) were prepd. by cyclizing EtO2CNHCSR with ClCH2CN and treating the thiazoles II with ClP(X)(OR1)2. I (R = OEt, R1 = Et, X = S) at 5 ppm gave 99% kill of Drosophila melanogaster in 1 hr. I also demonstrated acaricidal and nematocidal properties.
- IT 59965-62-7P 59965-65-0P 59965-72-9P 59965-80-9P

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. and condensation of, with chloroacetonitrile)

59965-62-7 CAPLUS RN

Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 3-cyclohexyl 1-ethyl ester CN (9CI) (CA INDEX NAME)

RN 59965-65-0 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 3-[(4-chlorophenyl)methyl] 1-ethyl ester (9CI) (CA INDEX NAME)

RN 59965-72-9 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 1-ethyl 3-(phenylmethyl) ester (9CI) (CA INDEX NAME)

RN 59965-80-9 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 1-ethyl 3-[(4-methylphenyl)methyl] ester (9CI) (CA INDEX NAME)

IT 59965-60-5P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)

(prepn. and reaction of, with potassium methylate)

RN 59965-60-5 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 3-butyl 1-ethyl ester, potassium salt (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} S & O \\ \parallel & \parallel \\ n\text{-BuO-C-NH-C-OEt} \end{array}$$

K

ANSWER 30 OF 39 CAPLUS COPYRIGHT 2002 ACS L6

1976:43357 CAPLUS AN

84:43357 DN

Alkyl S-aralkyl imidothiocarbonates TI

Takiguchi, Daigaku; Miyazaki, Koshin; Kato, Kinpei; Yasuda, Yasushi; IN Wakai, Akira

Nippon Soda Co., Ltd., Japan PA

Japan. Kokai, 9 pp. SO

CODEN: JKXXAF

DT Patent

Japanese LA

FAN.CNT 1

APPLICATION NO. DATE TP 50014601 PATENT NO. KIND DATE \_\_\_\_\_\_

JP 50014631 A2 19750215 JP 1973-66139 19730612 PI

ROC(O)N:C(SR1)(XR2) (I; R = lower alkyl; R1, R2 = lower alkyl, lower AΒ alkenyl, PhCH2, halobenzyl; X = O, S) were prepd. by treating ROC(O)NHC(S)XR1 (II) with R22SO4 or R2Y (Y = halo). I were effective components for fungicides. Thus, 17.1 g Et2SO4 was added to a mixt. of 28 ml 4N NaOH and 16.5 g II (R = Me, R1 = Et, X = S) below 10.degree. and the mixt. kept 2 hr at 30-5.degree. to give 15 g I (R = Me, R1 = R2 = Et, X = S). Among 20 more I prepd. were (R, R1, R2, X given): Et, Et, Et, O; Me, PhCH2, Et, S; Me, Me, Et, O; and Me, Me, Et, S.

51291-79-3P 57867-15-9P 57867-17-1P IT

57867-19-3P 57867-24-0P 57867-26-2P

57867-28-4P 57867-29-5P 57867-30-8P

57867-31-9P

RL: SPN (Synthetic preparation); PREP (Preparation)

(prepn. of)

51291-79-3 CAPLUS RN

Carbonimidothioic acid, (ethoxycarbonyl)-, O-ethyl S-methyl ester (9CI) CN (CA INDEX NAME)

57867-15-9 CAPLUS RN

Carbonimidothioic acid, (ethoxycarbonyl)-, diethyl ester (9CI) (CA INDEX CN NAME)

RN 57867-17-1 CAPLUS

CN Carbonimidothioic acid, (methoxycarbonyl)-, O-ethyl S-methyl ester (9CI) (CA INDEX NAME)

RN 57867-19-3 CAPLUS

CN Carbonimidothioic acid, (methoxycarbonyl)-, diethyl ester (9CI) (CA INDEX NAME)

RN 57867-24-0 CAPLUS

CN Carbonimidothioic acid, (methoxycarbonyl)-, S-[(3,4-dichlorophenyl)methyl] O-ethyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} \text{OEt} & \text{O} \\ \mid & \parallel \\ \text{C1} & \text{CH}_2\text{--} \text{s--} \text{C} \\ \end{array}$$

RN 57867-26-2 CAPLUS

CN Carbonimidothioic acid, (methoxycarbonyl)-, O-ethyl S-[(pentachlorophenyl)methyl] ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} \text{C1} & \text{OEt} & \text{O} \\ \text{C1} & \text{CH}_2\text{--} \text{s--} \text{C} \\ \text{C1} & \text{C1} \\ \end{array}$$

RN 57867-28-4 CAPLUS

CN Carbonimidothioic acid, (ethoxycarbonyl)-, S-ethyl O-2-propenyl ester

10/074,014

(9CI) (CA INDEX NAME)

$$\begin{array}{c|c} \text{O} & \text{SEt} \\ \parallel & \parallel & \parallel \\ \text{EtO-C-N} & \leftarrow \text{C-O-CH}_2\text{-CH} & \leftarrow \text{CH}_2 \end{array}$$

RN 57867-29-5 CAPLUS

CN Carbonimidothioic acid, [(1-methylethoxy)carbonyl]-, diethyl ester (9CI) (CA INDEX NAME)

RN 57867-30-8 CAPLUS

CN Carbonimidothioic acid, (ethoxycarbonyl)-, S-[(3,4-dichlorophenyl)methyl] O-ethyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} \text{OEt} & \text{O} \\ | & \text{|} \\ | & \text{|} \\ \text{C1} & \text{C-OEt} \\ \end{array}$$

RN 57867-31-9 CAPLUS

CN Carbonimidothioic acid, (ethoxycarbonyl)-, O-ethyl S-[(pentachlorophenyl)methyl] ester (9CI) (CA INDEX NAME)

C1 
$$CH_2-S-C=N-C-OEt$$

C1  $CH_2-S-C=N-C-OEt$ 

L6 ANSWER 31 OF 39 CAPLUS COPYRIGHT 2002 ACS

AN 1976:4965 CAPLUS

DN 84:4965

TI Insecticidal, acaricidal, and nematicidal O-triazolylthionophosphoric(phosphonic) acid esters or esteramides

IN Hoffmann, Hellmut; Hammann, Ingeborg; Homeyer, Bernhard; Stendel, Wilhelm

PA Bayer A.-G., Ger.

SO Ger. Offen., 45 pp. CODEN: GWXXBX

DT Patent LA German

FAN.CNT 1

two.	PATENT NO.	KIND	DATE	APPLICATION NO. DATE
ΡI	DE 2407304	A1	19750904	DE 1974-2407304 19740215
	SU 526275	D	19760825	SU 1975-2103867 19750211
	JP 50111231	A2	19750901	JP 1975-16963 19750212
	JP 50116476	A2	19750911	JP 1975-16962 19750212
	AT 7501061	Α	19750715	AT 1975-1061 19750213
	AT 329078	В	19760426	
	BE 825478	A1	19750813	BE 1975-153312 19750213
	SE 7501614	Α	19750821	SE 1975-1614 19750213
	DD 118514	С	19760312	DD 1975-184164 19750213
	PL 93295	P	19770530	PL 1975-178014 19750213
	NL 7501783	Α	19750819	NL 1975-1783 19750214
	FR 2261285	A1	19750912	FR 1975-4667 19750214
	DK 7500557	Α	19751013	DK 1975-557 19750214
	BR 7500909	Α	19751202	BR 1975-909 19750214
	ZA 7500943	Α	19760128	ZA 1975-943 19750214
	ES 434719	A1	19770201	ES 1975-434719 19750214
PRAI	DE 1974-2407304		19740215	

GI For diagram(s), see printed CA Issue.

AB Triazolyl phosphates I (R = R1 = OMe, OEt; R = OEt, R1 = NHCHMe2, NMe2, Ph; R2 = Me, allyl, CH2CN, CH2CH2CN, CH2CH:CHMe, CH2CMe:CH2; R3 = Me, Et, CHMe2) were prepd. by esterifying triazolols with RR1P(S)Cl. The triazolols were prepd. e.g. by cyclizing ethoxycarbonylthiosemicarbazides and alkylating the thiones. I are insecticides, acaricides, and nematocides. Thus I (R = R1 = OEt, R2 = CH2CN, R3 = Et) at 0.01% gave 100% kill of Phaedon cochleariae larva on cabbage leaves.

IT 5585-23-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP

(Preparation)

(prepn. and reaction of, with methylhydrazine)

RN 5585-23-9 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), diethyl ester (9CI) (CA INDEX NAME)

L6 ANSWER 32 OF 39 CAPLUS COPYRIGHT 2002 ACS

AN 1974:520638 CAPLUS

DN 81:120638

TI Acaricidal and insecticidal O-triazolyl phosphoro- and phosphonothioates

IN Hoffmann, Hellmut; Hammann, Ingeborg; Behrenz, Wolfgang; Homeyer, Bernhard; Stendel, Wilhelm

PA Bayer A.-G.

SO Ger. Offen., 51 pp.

CODEN: GWXXBX

DT Patent

LA German

FAN.CNT 1

PATENT NO. KIND DATE

APPLICATION NO. DATE

ΡI	DE 2301400	A1	19740718	DE	1973-2301400	19730112
11	DE 2301400 DE 2301400	C2	19841213	22	1575 2501100	13.00111
	HU 167093	P	19750828	ни	1974-BA3011	19740108
	NL 7400309	Ā	19740716		1974-309	19740109
	JP 49101544	A2	19740925		1974-5644	19740110
	JP 57036886	B4	19820806			
	DD 110164	C	19741212	DD	1974-175936	19740110
	AT 321944	В	19750425	ΑT	1974-180	19740110
	CS 175366	P	19770531	CS	1974-170	19740110
	CH 588504	Α	19770615	CH	1974-302	19740110
	JP 58038438	B4	19830823	JP	1974-5643	19740110
	BE 809633	A1	19740711	BE	1974-139713	19740111
	ZA 7400209	Α	19741127	ZA	1974-209	19740111
	AU 7464434	A1	19750717	AU	1974-64434	19740111
	GB 1406984	Α	19750924		1974-1371	19740111
	ES 422212	A1	19760501		1974-422212	19740111
	FR 2324640	A1	19770415		1974-1061	19740111
	SE 400769	С	19780720		1974-376	19740111
	CA 1050997	A1	19790320		1974-189956	19740111
	SU 713527	D	19800130		1974-1989776	19740111
	US 4229444	Α	19801021	US	1978-907388	19780518
PRAI	DE 1973-2301400		19730112			
	US 1974-430435		19740103		•	
CT	US 1976-645971		19760102			

GI For diagram(s), see printed CA Issue.

AB Sixteen phosphoro- and phosphonothioates I (R = OMe, SMe, SCH2CN, or SCH2CH:CH2; R1 = Me, Et, or CHMe2; R3 = Et, Ph, OMe, OEt, or NHCHMe2; R4 = Me, Et, or Pr) were prepd. in 58-88% yield by reaction of II with ClP(S)R3OR4 and used as acaricides and insecticides.

IT 5585-23-9P 51291-77-1P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP

(Preparation)

(prepn. and reaction with thiophosphorus acids)

RN 5585-23-9 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), diethyl ester (9CI) (CA INDEX NAME)

RN 51291-77-1 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 1-ethyl 3-methyl ester (9CI) (CA INDEX NAME)

L6 ANSWER 33 OF 39 CAPLUS COPYRIGHT 2002 ACS

AN 1974:82888 CAPLUS

DN 80:82888

- TI Purine studies. IX. Nucleophilic addition of barbituric acids to purines
- AU Pendergast, William
- CS Dep. Med. Chem., Aust. Natl. Univ., Canberra, Aust.
- SO J. Chem. Soc., Perkin Trans. 1 (1973), (22), 2759-63 CODEN: JCPRB4
- DT Journal
- LA English
- GI For diagram(s), see printed CA Issue.
- AB Purines with barbituric and 2-thiobarbituric acid underwent addn. reaction across the 1,6-double bond. E.g. 2-aminopurine with 2-thio-barbituric acid and 2,3-dihydropurin-2(3H)-one with barbituric acid gave 83% adduct (I) and 48% adduct (II) resp. The uv and NMR spectra of the adducts were reported and discussed.
- IT 51291-78-2P
- RN 51291-78-2 CAPLUS
- CN Carbonimidothioic acid, (ethoxycarbonyl)-, dimethyl ester (9CI) (CA INDEX NAME)

- L6 ANSWER 34 OF 39 CAPLUS COPYRIGHT 2002 ACS
- AN 1973:124479 CAPLUS
- DN 78:124479
- TI Organic sulfur compounds. X. Reactions of alkoxycarbonyl isothiocyanates with prim-.alpha.-acetylenic alcohol
- AU Nagano, Mitsuo; Matsui, Takashi; Tobitsuka, Junzo; Oyamada, Kozo
- CS Agric. Chem. Res. Lab., Sankyo Co., Ltd., Tokyo, Japan
- SO Chem. Pharm. Bull. (1973), 21(1), 62-73 CODEN: CPBTAL
- DT Journal
- LA English
- GI For diagram(s), see printed CA Issue.
- The reactions of alkoxycarbonyl isothiocyanates and .alpha.-acetylenic alcs. gave N-alkoxycarbonyl-O-acetylenyl thiocarbamates, N-alkoxycarbonyl-S allenyl thiolcarbamates and 4-alkylidene-2-alkoxycarbonylimino-1,3-oxathiolanes. The reaction patterns are dependent on the substituents on the 3-positions of .alpha.-acetylenic alcs. 3-Phenyl-2-propyn-1-ol react smoothly with RO2CNCS (I) to give 1:1 adducts which cyclize immediately to the 1,3-oxathiolanes (II, R = Me, Et Bu, etc.; R1 = Ph). 2-Butyn-1-ol reacts slowly with I to give N-alkoxtcarbonyl-O-2- butynyl thiocarbamates, which cyclize to 4-ethylidene-1,3-oxathiolanes (II) (R = Me, Et, Bu, etc.; R1 Me) even in the presence of a base. In the case of 2- propyn-1-ol, HC:CCH2O-CSNHCO2R, H2C:- C:CHSCONHCO2R and II (R = Me, Et, Pr, etc.; R1 = H) are obtained. The cyclization mechanisms were detd. by using HC:CCH2OCSND- CP2CHMe2.
- IT 37063-41-5P 37063-42-6P 37063-43-7P 37063-44-8P 37063-45-9P 40914-41-8P 40914-47-4P 40914-50-9P 40914-51-0P 40914-54-3P 40914-56-5P 40914-57-6P 40914-58-7P 40914-72-5P 40942-44-7P RL: SPN (Synthetic preparation); PREP (Preparation)

(prepn. of)

RN 37063-41-5 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 1-methyl 3-(2-propynyl) ester (9CI) (CA INDEX NAME)

RN 37063-42-6 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 1-ethyl 3-(2-propynyl) ester (9CI) (CA INDEX NAME)

RN 37063-43-7 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 1-propyl 3-(2-propynyl) ester (9CI) (CA INDEX NAME)

RN 37063-44-8 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 3-(2-butynyl) 1-methyl ester (9CI) (CA INDEX NAME)

RN 37063-45-9 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 1-ethyl 3-(3-phenyl-2-propynyl) ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & \text{O} & \text{S} \\ || & || \\ \text{EtO-C-NH-C-O-CH}_2\text{-C} \end{array} \text{C-Ph}$$

RN 40914-41-8 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 1-ethyl 3-(2-oxopropyl) ester (9CI) (CA INDEX NAME)

RN 40914-47-4 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 1-(1-methylethyl) 3-(2-propynyl) ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} \text{O} & \text{S} \\ \parallel & \parallel \\ \text{i-PrO-C-NH-C-O-CH}_2\text{-C} \end{array} \text{CH}$$

RN 40914-50-9 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 1-butyl 3-(2-propynyl) ester (9CI) (CA INDEX NAME)

RN 40914-51-0 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 1-(2-methylpropyl) 3-(2-propynyl) ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} O & S \\ \parallel & \parallel \\ i-BuO-C-NH-C-O-CH_2-C \Longrightarrow CH \end{array}$$

RN 40914-54-3 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 3-(2-butynyl) 1-ethyl ester (9CI) (CA INDEX NAME)

RN 40914-56-5 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 3-(2-butynyl) 1-(1-methylethyl) ester (9CI) (CA INDEX NAME)

RN 40914-57-6 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 1-butyl 3-(2-butynyl) ester

10/074,014

(9CI) (CA INDEX NAME)

RN 40914-58-7 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 3-(2-butynyl) 1-(2-methylpropyl) ester (9CI) (CA INDEX NAME)

RN 40914-72-5 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NDC(S)(OH)), 1-(1-methylethyl) 3-(2-propynyl) ester (9CI) (CA INDEX NAME)

RN 40942-44-7 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 3-(2-butynyl) 1-propyl ester (9CI) (CA INDEX NAME)

L6 ANSWER 35 OF 39 CAPLUS COPYRIGHT 2002 ACS

AN 1973:97628 CAPLUS

DN 78:97628

TI Insecticidal thiazolo(thiono)phosphoric(or phosphonic) acid esters

PA Farbenfabriken Bayer A.-G.

SO Fr., 29 pp.

CODEN: FRXXAK

DT Patent

LA French

FAN.CNT 2

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
ΡI	FR 2121065	A5	19720818	FR 1971-47274	19711229
	DE 2064307	Α	19720706	DE 1970-2064307	19701229
PRAI	DE 1970-2064307		19701229		

GI For diagram(s), see printed CA Issue.

AB Thiazolylphosphoric esters I (R = EtO, Et; X = O, S; R1 = Me, Et, Me2CH; R2 = Et, Me2CH) were prepd. in 59-85% yield by treating R(EtO)P(X)Cl with the appropriate 4-hydroxythiazole. O-Isopropyl-O-(2-ethoxy-5-

isopropoxycarbonylthiazol-4-yl) methylthiophosphonate was similarly prepd. The 4-hydroxythiazoles were obtained by cyclizing all Eto2CN:C(OR1)SCH2CO2R2.

IT 40509-96-4P

RN 40509-96-4 CAPLUS

CN Acetic acid, [[[(ethoxycarbonyl)imino](1-methylethoxy)methyl]thio]-, 1-methylethyl ester (9CI) (CA INDEX NAME)

- L6 ANSWER 36 OF 39 CAPLUS COPYRIGHT 2002 ACS
- AN 1973:58366 CAPLUS

DN 78:58366

- TI Organic sulfur compounds. VIII. Reaction of alkoxycarbonyl isothiocyanates and 2-aminothiazole
- AU Nagano, Mitsuo; Tobitsuka, Junzo; Matsui, Takashi; Oyamada, Kozo
- CS Agric. Chem. Res. Lab., Sankyo Co., Ltd., Tokyo, Japan
- SO Chem. Pharm. Bull. (1972), 20(12), 2618-25 CODEN: CPBTAL
- DT Journal
- LA English
- GI For diagram(s), see printed CA Issue.
- AB The reactions of some alkoxycarbonyl isothiocyanates with 2-aminothiazole (II) afforded thiazolo[3,2-a]-s-triazine-4-thion-2-one (I), N-alkoxycarbonyl-N'-(2-thiazolyl)thioureas, Alkyl-N-(2-thiazolyl)-carbamates, N-alkoxycarbonyl thiocarbamates and HSCN. However, in the case using PhO2CNCS (III), the corresponding 1:1 adduct of II and III could not be obtained, but thiazolo[3,2-a]-s-thiazine-2-thion-4-one (IV) was isolated, besides I, phenyl 2-thiazolylcarbamate, and phenol.
- IT 39142-28-4P 39142-31-9P 39142-33-1P

39142-36-4P 39142-39-7P

RN 39142-28-4 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), dimethyl ester (9CI) (CA INDEX NAME)

RN 39142-31-9 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), dipropyl ester (9CI) (CA INDEX NAME)

RN 39142-33-1 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), bis(1-methylethyl) ester (9CI) (CA INDEX NAME)

RN 39142-36-4 CAPLUS

$$\begin{array}{c|c} O & S \\ \parallel & \parallel \\ n\text{-BuO-} C\text{--} NH\text{--} C\text{--} OBu\text{--} n \end{array}$$

RN 39142-39-7 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), bis(2-methylpropyl) ester (9CI) (CA INDEX NAME)

L6 ANSWER 37 OF 39 CAPLUS COPYRIGHT 2002 ACS

AN 1973:58365 CAPLUS

DN 78:58365

TI Organic sulfur compounds. IX. Reaction of ethoxycarbonyl isothiocyanate with 4,5-substituted 2-aminothiazoles

AU Nagano, Mitsuo; Matsui, Takashi; Tobitsuka, Junzo; Oyamada, Kozo

CS Agric. Chem. Res. Lab., Sankyo Co., Ltd., Tokyo, Japan

SO Chem. Pharm. Bull. (1972), 20(12), 2626-33 CODEN: CPBTAL

DT Journal

LA English

GI For diagram(s), see printed CA Issue.

The Reactions of SCNCO2Et with 4,5-substituted 2-aminothiazoles afforded thiazolo[3,2-a]-s-triazine-4-thion-2-ones (I), N-alkoxycarbonyl-N'-(2-thiazolyl)thioureas (II), alkyl-N-(2-thiazolyl)carbamates (III) (R = H, Me, Ph; R1 = H, Me, Et, Pr, Bu), EtO2CNHC(:S)OEt, and HSCN. However, in the cases of the amines whose pKa values were smaller than that of 2-aminothiazole or the amines which had some substituents on the 4-position the corresponding cyclic compds. (25) could not be obtained. A series of these phenomena was discussed in connection with the basicities of the 2-aminothiazoles and the steric hindrance of the substituents on

the 4-position.

IT 5585-23-9P

RL: SPN (Synthetic preparation); PREP (Preparation)

(prepn. of)

RN 5585-23-9 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), diethyl ester (9CI) (CA INDEX NAME)

L6 ANSWER 38 OF 39 CAPLUS COPYRIGHT 2002 ACS

AN 1972:526044 CAPLUS

DN 77:126044

TI Carbamates

IN Oyamada, Kozo; Nagano, Mitsuo; Tobizuka, Junzo; Matsui, Takashi; Saito, Masataka

PA Sankyo Co., Ltd.

SO Jpn. Kokai Tokkyo Koho, 5 pp.

CODEN: JKXXAF

DT Patent

LA Japanese

FAN.CNT 1

PATENT NO. KIND DATE APPLICATION NO. DATE

AB Carbamates, RC.tplbond.CCH2OC(:S)NHC(:O)OR1 (I), useful as insecticides, analgesics, and antiinflammatories, were prepd. by the reaction of acetylene alc., RC.tplbond.CCH2OH (II), with isothiocyanate, SCNC(:O)OR1 (III). Thus, a mixt. of 1.12 g II (R = H) and 2.34 g III (R1 = Me) in AcOEt was stirred 5 hr to give 0.82 g I (R = H, R1 = Me). Among 14 more I similarly prepd. were (R and R1 given): H, Et; H, Pr; Me, Me; Ph, Et; Me, Ph

IT 37063-41-5P 37063-42-6P 37063-43-7P 37063-44-8P 37063-45-9P 37063-46-0P

RL: SPN (Synthetic preparation); PREP (Preparation)

(prepn. of)

RN 37063-41-5 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 1-methyl 3-(2-propynyl) ester (9CI) (CA INDEX NAME)

RN 37063-42-6 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 1-ethyl 3-(2-propynyl) ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} o & s \\ || & || \\ \text{Eto-C-NH-C-O-CH}_2\text{-C} = \text{CH} \end{array}$$

RN 37063-43-7 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 1-propyl 3-(2-propynyl) ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & \text{O} & \text{S} \\ || & || \\ \\ \text{n-Pro-C-NH-C-O-CH}_2\text{-C} \end{array} \text{CH}$$

RN 37063-44-8 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 3-(2-butynyl) 1-methyl ester (9CI) (CA INDEX NAME)

RN 37063-45-9 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 1-ethyl 3-(3-phenyl-2-propynyl) ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} O & S \\ \parallel & \parallel \\ EtO-C-NH-C-O-CH_2-C \Longrightarrow C-Ph \end{array}$$

RN 37063-46-0 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 3-(2-butynyl) 1-phenyl ester (9CI) (CA INDEX NAME)

L6 ANSWER 39 OF 39 CAPLUS COPYRIGHT 2002 ACS

AN 1972:33982 CAPLUS

DN 76:33982

TI N-Acylcarbothioamides

IN Grigat, Ernst

PA Farbenfabriken Bayer A.-G.

SO Ger. Offen., 20 pp.

CODEN: GWXXBX

DT Patent

LA German

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE	
ΡI	DE 2017966	Α	19711028	DE 1970-2017966	19700415	
AB	Thirteen title c	ompds.	, RC(S)NHCO	R1 (I, e.g. R=Cl3CCH2	O, p-02NC6H4O,	
morpholino, Me2N, or PhO, R1=Et, CCl3, C6H4Cl-p, OCH2CH2Cl, Ph,						
	CH2OC6H3Cl2-2,4,	or $4,$	5,6-trichlo	ro-2-pyrimidinyl), we	re prepd. by	
	reaction of RClC	:NCOR1	with H2S o	r H2S-releasing compd	s. Thus,	
	CC13CH2OCC1:NCOE	t in E	t20 was add	ed to Et3N in Et2O sa	td. with H2S at	
	0.degree. to giv	e 83%	I (R=Cl3CCH	20, R1=Et). Similarl	y prepd. were 12	
	other I.					

IT 34840-04-5P 34840-54-5P

RN 34840-04-5 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 1-(4-chlorophenyl) 3-phenyl ester (9CI) (CA INDEX NAME)

RN 34840-54-5 CAPLUS

CN Thioimidodicarbonic acid ((HO)C(O)NHC(S)(OH)), 1-(2-chloroethyl) 3-(2,4-dimethylphenyl) ester (9CI) (CA INDEX NAME)

=> log y SINCE FILE TOTAL COST IN U.S. DOLLARS ENTRY SESSION 314.87 FULL ESTIMATED COST 174.38 DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE TOTAL ENTRY SESSION -24.16 -24.16CA SUBSCRIBER PRICE

STN INTERNATIONAL LOGOFF AT 11:10:53 ON 03 OCT 2002

1

K

L6 ANSWER 30 OF 39 CAPLUS COPYRIGHT 2002 ACS

AN 1976:43357 CAPLUS

DN 84:43357

TI Alkyl S-aralkyl imidothiocarbonates

IN Takiguchi, Daigaku; Miyazaki, Koshin; Kato, Kinpei; Yasuda, Yasushi; Wakai, Akira

PA Nippon Soda Co., Ltd., Japan

SO Japan. Kokai, 9 pp. CODEN: JKXXAF

DT Patent

LA Japanese

FAN.CNT 1

PATENT NO. KIND DATE APPLICATION NO. DATE

PI JP 50014631 A2 19750215 JP 1973-66139 19730612

AB ROC(O)N:C(SR1)(XR2) (I; R = lower alkyl; R1, R2 = lower alkyl, lower alkenyl, PhCH2, halobenzyl; X = O, S) were prepd. by treating ROC(O)NHC(S)XR1 (II) with R22SO4 or R2Y (Y = halo). I were effective components for fungicides. Thus, 17.1 g Et2SO4 was added to a mixt. of 28 ml 4N NaOH and 16.5 g II (R = Me, R1 = Et, X = S) below 10.degree. and the mixt. kept 2 hr at 30-5.degree. to give 15 g I (R = Me, R1 = R2 = Et, X = S). Among 20 more I prepd. were (R, R1, R2, X given): Et, Et, Et, O; Me, PhCH2, Et, S; Me, Me, Et, O; and Me, Me, Et, S.

IT 51291-79-3P 57867-15-9P 57867-17-1P 57867-19-3P 57867-24-0P 57867-26-2P 57867-28-4P 57867-29-5P 57867-30-8P

57867-31-9P

RL: SPN (Synthetic preparation); PREP (Preparation)

(prepn. of) 51291-79-3 CAPLUS

CN Carbonimidothioic acid, (ethoxycarbonyl)-, O-ethyl S-methyl ester (9CI) (CA INDEX NAME)

RN

RN 57867-15-9 CAPLUS

CN Carbonimidothioic acid, (ethoxycarbonyl)-, diethyl ester (9CI) (CA INDEX NAME)

RN 57867-17-1 CAPLUS

SMe O | | | | EtO-C== N-C-OMe

RN 57867-19-3 CAPLUS

CN Carbonimidothioic acid, (methoxycarbonyl)-, diethyl ester (9CI) (CA INDEX NAME)

RN 57867-24-0 CAPLUS

CN Carbonimidothioic acid, (methoxycarbonyl)-, S-[(3,4-dichlorophenyl)methyl] O-ethyl ester (9CI) (CA INDEX NAME)

C1 C1 CH2-S-C=N-C-OME

RN 57867-26-2 CAPLUS

CN Carbonimidothioic acid, (methoxycarbonyl)-, O-ethyl S-[(pentachlorophenyl)methyl] ester (9CI) (CA INDEX NAME)

 $\begin{array}{c|c} \text{C1} & \text{OEt} & \text{O} \\ \mid & \parallel & \parallel \\ \text{C1} & \text{C1} & \text{C1} \\ \end{array}$ 

RN 57867-28-4 CAPLUS

CN Carbonimidothioic acid, (ethoxycarbonyl)-, S-ethyl O-2-propenyl ester

10/074,014

(9CI) (CA INDEX NAME)

$$\begin{array}{c|c} \text{O} & \text{SEt} \\ \parallel & \parallel \\ \text{EtO-C-N----} \text{C-O-CH}_2\text{--CH-----} \text{CH}_2 \end{array}$$

RN 57867-29-5 CAPLUS

CN Carbonimidothioic acid, [(1-methylethoxy)carbonyl]-, diethyl ester (9CI) (CA INDEX NAME)

RN 57867-30-8 CAPLUS

CN Carbonimidothioic acid, (ethoxycarbonyl)-, S-[(3,4-dichlorophenyl)methyl] O-ethyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} \text{OEt} & \text{O} \\ & \parallel \\ \text{CH}_2 - \text{S} - \text{C} & \text{N} - \text{C} - \text{OEt} \\ \end{array}$$

RN 57867-31-9 CAPLUS

CN Carbonimidothioic acid, (ethoxycarbonyl)-, O-ethyl S- [(pentachlorophenyl)methyl] ester (9CI) (CA INDEX NAME)

C1 
$$CH_2-S-C=N-C-OEt$$

C1  $CI$   $CI$   $CI$   $CI$ 

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AN 1976:4965 CAPLUS

DN 84:4965

TI Insecticidal, acaricidal, and nematicidal O-triazolylthionophosphoric(phosphonic) acid esters or esteramides

IN Hoffmann, Hellmut; Hammann, Ingeborg; Homeyer, Bernhard; Stendel, Wilhelm

PA Bayer A.-G., Ger.

SO Ger. Offen., 45 pp. CODEN: GWXXBX